Section I (Amendments to the Claims)

Please amend claims 1, 3, 4, 6, 7, 9, 10, 12, 13, 15, and 46-48, as set out in the following listing of the claims of the application.

Please cancel claims 2, 5, 8, 14, and 62 without prejudice.

1. (Currently amended) A nucleic acid construct encoding a fusion protein, wherein the construct comprises a coding sequence for a protein of interest and a coding sequence for a protease <u>subtilisin</u> prodomain protein, wherein the fusion protein comprises the protein of interest operatively linked to the <u>protease subtilisin</u> prodomain protein, <u>and</u> wherein the <u>protease subtilisin</u> prodomain protein <u>has binding with high affinity binds</u> to a <u>subtilisin-protease</u> or a variant thereof <u>with a Kd of 10 nM or less</u>, and wherein the subtilisin variant retains the activity of subtilisin.

2. (Cancelled)

- 3. (Currently amended) The nucleic acid construct according to claim [[2]]], wherein the protease subtilisin prodomain protein further comprises one or more amino acid substitutions that increase binding affinity for subtilisin or a variant thereof, as compared to the protease subtilisin prodomain protein with no substitutions.
- 4. (Currently amended) The nucleic acid construct according to claim 1, wherein the protease subtilisin prodomain protein comprises a variant of SEQ ID NO: 1, wherein the variant comprises a substitution at one or more of positions P1-P4 wherein the substitution comprises any of F or Y substituted for P4, any amino acid residue substituted for P3, A or S substituted for P2 and M, F, Y H, or L substituted for P1.

- 6. (Currently amended) The nucleic acid construct according to claim [[5]]L, wherein the protease <u>subtilisin</u> prodomain protein comprises substitutions of amino acid residues F or Y for P4, any amino acid residue for P3, A or S for P2 and M, F, Y, H, or L for P1 at the C-terminal end.
- (Currently amended) A fusion protein comprising a target protein operatively linked to a protease subtilisin prodomain protein, wherein the protease subtilisin prodomain protein is

modified to exhibit an increased affinity for subtilisin or a variant thereof, as compared to the unmodified protease <u>subtilisin</u> prodomain protein, <u>and wherein the subtilisin variant retains the</u> activity of subtilisin.

- (Currently amended) The fusion protein according to claim [[8]]Z, wherein the protease subtilisin prodomain protein comprises substitution of amino acids P4-P1with the amino acid sequence FKAM.
- (Currently amended) The fusion protein according to claim 7, wherein the protease <u>subtilisin</u> prodomain protein comprises the amino acid sequence E E D K L (F/Y) Q S (M/L/Y) (SEQ ID NO: 7).
- 11. (Previously Presented) The fusion protein according to claim 7, wherein the target protein is staphylococcal Protein AB domain; Protein AB mutant A219; Streptococcal protein GB domain; Streptococcal protein Ga domain; Protein GB mutant G311; E. coli hypothetical Yab; Bovine a-subunit of transducin; M. thermautotrophicus CDC6; streptavidin; avidin; Taq polymerase; alkaline phosphatase; RNase; DNase; restriction enzymes; peroxidases; endo-1, 4-beta glucanase; endo-1, 3-beta-glucanase; chitinases; beta and alpha glucosidases; beta and alpha glucoronidases; amylase; glucosyl-transferases; phospho-transferases; chloramphenicol-acetyl-transferase; beta-lactamase; luciferase; esterases; lipases; proteases; bacteriocines; antibiotics; enzyme inhibitors; growth factors; hormones; receptors; membrane proteins; nuclear proteins; transcriptional factors; translational factors or nucleic acid modifyine enzymes.
- 12. (Currently amended) A DNA construct for the preparation of a fusion protein, wherein the construct comprises a coding sequence of a protein of interest and a DNA sequence encoding a subtilisin binding protein having binding with high affinity for which binds to subtilisin with a Kd of 10 nM or less.
- 13. (Currently amended) A method for the production of a subtilisin binding fusion protein, the method comprising: providing a nucleic acid construct encoding a fusion protein wherein the fusion protein comprises a protease subtilisin prodomain protein operatively linked to a second protein of interest, wherein the protease subtilisin prodomain protein is modified to bind subtilisin or a variant thereof with increased affinity as compared to an unmodified protease

subtilisin prodomain protein, and wherein the subtilisin variant retains the activity of subtilisin; transfecting a host cell with the nucleic acid construct; and culturing the transformed host cell under conditions suitable for expression of the fusion protein.

- (Currently amended) The method according to claim [[14]]13, wherein the subtilisin prodomain is modified by replacing the P4 through P1 amino acids with FKAM, FKAY or FKAF.
- 16. (Previously Presented) The method according to claim 15, wherein the second protein of interest is staphylococcal Protein AB domain; Protein AB mutant A219; Streptococcal protein GB domain; Streptococcal protein Ga domain; Brotein GB mutant G311; E. coli hypothetical Yab; Bovine a-subunit of transducin; M. thermautotrophicus CDC6; streptavidin; avidin; Taq polymerase; alkaline phosphatase; RNase; DNase; restriction enzymes; peroxidases; endo-1, 4-beta glucanase; endo-1, 3-beta-glucanase; chitinases; beta and alpha glucosidases; beta and alpha glucoronidases; amylase; glucosyl-transferases; phospho-transferases; chloramphenicol-acetyl-transferase; beta-lactamase; luciferase; esterases; lipases; proteases; bacteriocines; antibiotics; enzyme inhibitors; growth factors; hormones; receptors; membrane proteins; nuclear proteins; transcriptional factors; translational factors or nucleic acid modifying enzymes.
- 17. (Original) The method according to claim 13, wherein the host cells includes cells from, Escherichia coli, Bacillus, Salmonella, Pseudomonas; Saccharomyces cerevisiae, Pichia pastoris, Kluveromyces, Candida, Schizosaccharomyces; or CHO cells.
- 18. (Withdrawn) A method for purifying a protein of interest from a fusion protein and separation therefrom, the method comprising: contacting a fusion protein comprising a protease prodomain protein operatively linked to the protein of interest with an effective amount of subtilisin or a variant thereof under conditions suitable for the formation of a binding complex between the subtilisin or variant thereof and the protease prodomain protein of the fusion protein; incubating the binding complex for a sufficient time for the subtilisin or variant thereof to cleave the protein of interest from the binding complex; and recovering the protein of interest.
- 19. (Withdrawn) The method according to claim 18, wherein the subtilisin has been modified to specifically bind to the protease prodomain fusion protein.

- (Withdrawn) The method according to claim 19, wherein the subtilisin includes mutations Q2K, S3C, P5S, K43N, A73L, deletion of 75-83, E156S, G166S, G169A, S188P, Q206C, N212G, K217L, N218S, T254A, Q271E, Y104A, G128S and at least one additional mutation at amino acid position 32, 155 or 221.
- 21. (Withdrawn) The method according to claim 19, wherein the protease prodomain protein is a subtilisin prodomain and modified by replacing the P4 through P1 amino acids with FKAM, FKAY or FKAF
- 22. (Withdrawn) The method according to claim 21, wherein the protein of interest is staphylococcal Protein AB domain; Protein AB mutant A219; Streptococcal protein GB domain; Streptococcal protein Ga domain; Protein GB mutant G311; E. coli hypothetical Yab; Bovine a-subunit of transducin; M. thermautotrophicus CDC6; streptavidin; avidin; Taq polymerase; alkaline phosphatase; RNase; DNase; restriction enzymes; peroxidases; endo-1,4-beta glucanase; endo-1, 3-beta-glucanase; chitinases; beta and alpha glucoronidases; amylase; glucosyl-transferases; phospho-transferases; chloramphenicol-acetyl-transferase; beta-lactamase; luciferase; esterases; lipases; proteases; bacteriocines; antibiotics; enzyme inhibitors; growth factors; translational factors or nucleic acid modifying enzymes.
- 23. (Withdrawn) The method according to claim 20, wherein the subtilisin is immobilized on a solid phase matrix.
- 24. (Withdrawn) The method according to claim 21, wherein the prodomain of subtilisin is mutated to increase binding affinity of subtilisin to greater than $109 \, M^{-1}$.
- 25. (Withdrawn) The method according to claim 19, wherein the subtilisin includes mutations Q2K, S3C, P5S, K43N, A73L, deletion of 75-83, E156S, G169A, S188P, Q206C, N212G, K217L, N218S, T254A, Q271E, Y104A, G128S and at least one additional mutation at amino acid position 32, or 221.
- (Withdrawn) The method according to claim 20, wherein the subtilisin is S189, S190, S194, S196, S197, or S198.

- (Withdrawn) The method according to claim 25, wherein the subtilisin is \$199, \$201 or
- 28. (Withdrawn) An assay method for detecting the presence of a substance of interest in a test sample comprising: (a) incubating a test sample, which may contain a substance of interest, with a sufficient amount of a protease prodomain fusion protein, wherein the protease prodomain fusion protein comprises: (i) a protease prodomain capable of binding with high affinity to a subtilisin or variant thereof, and (ii) a second protein capable of binding the substance of interest, under incubating conditions that allow for the binding of the substance of interest to the second protein; (b) contacting the protease prodomain fusion protein used in step (a) to subtilisin or a variant thereof, wherein the subtilisin or a variant thereof is in solution in an amount effective to bind the fusion protein or immobilized on a solid phase to form a subtilisin/prodomain fusion protein binding complex; (c) incubating the subtilisin/prodomain fusion protein binding complex; (c) incubating the subtilisin/prodomain fusion protein binding complex; (c) recovering the second protein bound to the substance of interest.
- 29. (Withdrawn) The method according to claim 28, further comprising introducing a detectable label capable of binding to the substance of interest; and determining the presence or absence of the label, to provide an indication of the presence or absence of the substance of interest in the test sample.
- 30. (Withdrawn) The method according to claim 29, wherein the detectable label is introduced before separation of the second protein from the binding complex or after the second protein is recovered.
- 31. (Withdrawn) The method according to claim 28, wherein the test sample is blood, urine, semen, saliva, mucus, tears, or vaginal secretions.
- 32. (Withdrawn) The method according to claim 31, wherein the substance of interest is an antibody.
- 33. (Withdrawn) The method according to claim 32, wherein the second protein is an antigenic receptor having affinity for the antibody.

- 34. (Withdrawn) The method according to claim 31, wherein the substance of interest is an antieen.
- 35. (Withdrawn) The method according to claim 34, wherein the second protein is an antibody having affinity for the antibody.
- 36. (Withdrawn) The method according to claim 28, wherein the subtilisin has been modified to specifically bind to the protease prodomain fusion protein.
- 37. (Withdrawn) The method according to claim 36, wherein the subtilisin includes mutations Q2K, S3C, P5S, K43N, A73L, deletion of 75-83, E156S, G166S, G169A, S188P, Q206C, N212G, K217L, N218S, T254A, Q271E, Y104A, G128S and at least one additional mutation at amino acid position 32, 155 or 221.
- 38. (Withdrawn) The method according to claim 28, wherein the protease prodomain protein is a subtilisin prodomain and modified by replacing the P4 through P1 amino acids with FKAM, FKAY or FKAF.
- 39. (Withdrawn) A drug delivery system comprising a subtilisin prodomain protein associated with a drug of interest to form a fusion product, wherein the fusion product is further complexed to a subtilisin or variant thereof to form a drug delivery complex.
- 40. (Withdrawn) The drug delivery system according to claim 39, wherein the drug of interest is conjugated to the subtilisin prodomain protein either directly or through a linker moiety.
- 41. (Withdrawn) The drug delivery system according to claim 39, wherein the drug of interest is slowly released from the drug delivery complex.
- 42. (Withdrawn) The drug delivery system according to claim 41, wherein the drug delivery product is included in a composition and administered parenterally, orally, topically or by inhalation
- (Withdrawn) The drug delivery system according to claim 41, wherein the composition comprises a solid, gel, liquid or aerosol.

- 44. (Withdrawn) The drug delivery system according to claim 41, wherein the subtilisin includes mutations Q2K, S3C, P5S, K43N, A73L, deletion of 75-83, E156S, G166S, G169A, S188P, Q206C, N212G, K217L, N218S, T254A, Q271E, Y104A, G128S and at least one additional mutation at amino acid position 32, 155 or 221.
- 45. (Withdrawn) The drug delivery system according to claim 41, wherein the subtilisin prodomain protein is modified by replacing the P4 through P1 amino acid residues with FKAM, FKAY or FKAF.
- 46. (Currently amended) A nucleic acid construct encoding a fusion protein, wherein the construct comprises a coding sequence for a protein of interest and a coding sequence for a second protein, wherein the second protein binds with high affinity to a <u>subtilisin-protease</u> or a variant thereof with a Kd of 10 nM or less, and wherein the subtilisin variant retains the activity of subtilisin.
- 47. (Currently amended) A nucleic acid construct according to claim 46, wherein the fusion protein comprises the protein of interest linked to the second protein by a peptide bond and wherein the <u>protease subtilisin</u> hydrolyzes the peptide bond.
- 48. (Currently amended) A nucleic acid construct according to claim 46, wherein P1, P2 and P4 amino acids of the second protein generate affinity for S1, S2 and S4 binding pockets of the subtilisin-protease-or a variant thereof.
- 49. (Previously Presented) A nucleic acid construct according to claim 48, wherein the second protein comprises amino acid residues F or Y at the P4 position, any amino acid residue at the P3 position. A. S. V. or T at the P2 position and M. F. Y. H. or L at the P1 position.
- 50. (Withdrawn) A protease variant that is altered to specifically hydrolyze a fusion protein upon addition of a chemical trigger and the fusion protein comprises a binding sequence for a protease fused to a protein of interest.
- 51. (Withdrawn) A protease variant according to claim 50, wherein the altered protease is a subtilisin variant.

- 52. (Withdrawn) A protease variant according to claim 51, wherein the subtilisin variant comprises a mutation at amino acid 32.
- 53. (Withdrawn) A method of producing a protein of interest, comprising generating a fusion protein comprising a binding sequence for a protease fused to a protein of interest, and reacting said fusion protein with a protease variant that is altered to specifically hydrolyze the fusion protein and yield said protein of interest upon addition of a chemical trigger, wherein the reaction is conducted in the presence of said chemical trigger, and recovering said protein of interest.
- 54. (Withdrawn) A nucleic acid construct encoding a fusion protein, wherein the construct comprises a coding sequence for a protein of interest that is operatively linked to coding sequence for a peptide, wherein the peptide generates affinity for a protease or a variant thereof.
- 55. (Withdrawn) A nucleic acid construct according to claim 54, wherein the protease hydrolyzes a peptide bond joining the peptide to the protein of interest.
- 56. (Withdrawn) A nucleic acid construct according to claim 54, wherein P1, P2 and P4 amino acids of the peptide generate affinity for S1, S2 and S4 binding pockets of protease or a variant thereof.
- 57. (Withdrawn) A nucleic acid construct according to claim 56, wherein the peptide comprises amino acid residues F or Y at the P4 position, any amino acid residue at the P3 position, A, S, V, or T at the P2 position and M, F, Y, H, or L at the P1 position.
- 58. (Withdrawn) A method for the production of a subtilisin binding fusion protein, the method comprising: providing a nucleic acid construct encoding a fusion protein wherein the fusion protein comprises a peptide and a second protein of interest, wherein the peptide is modified to bind subtilisin or a variant thereof with high affinity; transfecting a host cell with the nucleic acid construct; and culturing the transformed host cell under conditions suitable for expression of the fusion protein.
- 59. (Withdrawn) A method for purifying a protein of interest from a fusion protein and separation therefrom, the method comprising: contacting a fusion protein comprising a peptide operatively linked to the protein of interest with an effective amount of subtilisin or a variant thereof under conditions suitable for the formation of a binding complex between the subtilisin

or variant thereof and the peptide of the fusion protein; incubating the binding complex for a sufficient time for the subtilisin or variant thereof to cleave the protein of interest from the binding complex; and recovering the protein of interest.

60. (Withdrawn) An assay method for detecting the presence of a substance of interest in a test sample comprising: (a) incubating a test sample, which may contain a substance of interest, with a sufficient amount of a fusion protein comprising: (i) a peptide capable of binding with high affinity to a subtilisin or variant thereof, and (ii) a second protein capable of binding the substance of interest, under incubating conditions that allow for the binding of the substance of interest to the second protein; (b) contacting the fusion protein used in step (a) to subtilisin or a variant thereof, wherein the subtilisin or a variant thereof is in solution in an amount effective to bind the fusion protein or immobilized on a solid phase to form a subtilisin/fusion protein binding complex; (c) incubating the subtilisin/fusion protein binding complex for a sufficient time for the subtilisin or variant thereof to cleave the second protein from the binding complex; (d) recovering the second protein bound to the substance of interest.

61. (Withdrawn) A drug delivery system comprising a peptide generating affinity for a subtilisin or a variant thereof associated with a drug of interest to form a fusion product, wherein the fusion product is further complexed to said subtilisin or variant thereof to form a drug delivery complex.